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Carslaw, Nicola orcid.org/0000-0002-5290-4779 (2017) Observations and impacts of bleach washing on indoor chlorine chemistry. Indoor air. pp. 1082-1090. ISSN 0905-6947

<https://doi.org/10.1111/ina.12402>

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Observations and impacts of bleach washing on indoor chlorine chemistry

J.P.S Wong,^a N. Carslaw,^b R. Zhao,^c S. Zhou,^d and J.P.D. Abbatt^{d,*}

*to whom correspondence should be addressed

^a School of Earth and Atmospheric Sciences, Georgia Institute of Technology, 311 Ferst Drive, Atlanta, GA 30331 USA

^b Environment Department, University of York, Wentworth Way, Heslington, York YO10 5NG, UK

^c Division of Chemistry and Chemical Engineering, California Institute of Technology, 1200 E California Blvd, Pasadena, CA 91125 USA

^d Department of Chemistry, University of Toronto, 80 St. George Street, Toronto, ON M5S 3H6 Canada

Abstract

Ambient levels of chlorinated gases and aerosol components were measured by on-line chemical ionization and aerosol mass spectrometers after an indoor floor was repeatedly washed with a commercial bleach solution. Gaseous chlorine (Cl_2 , 10's of ppbv) and hypochlorous acid (HOCl , 100's of ppbv) arise after floor washing, along with nitryl chloride (ClNO_2), dichlorine monoxide (Cl_2O) and chloramines (NHCl_2 , NCl_3). Much higher mixing ratios would prevail in a room with lower and more commonly encountered air exchange rates than that observed in the study (12.7 h^{-1}). Coincident with the formation of gas-phase species, particulate chlorine levels also rise. Cl_2 , ClNO_2 , NHCl_2 and NCl_3 exist in the headspace of the bleach solution whereas HOCl was only observed after floor washing. HOCl decays away 1.4 times faster than the air exchange rate, indicative of uptake onto room surfaces and consistent with the well-known chlorinating ability of HOCl . Photochemical box modeling captures the temporal profiles of Cl_2 and HOCl very well, and indicates that the OH , Cl and ClO gas-phase radical concentrations in the indoor environment could be greatly enhanced ($> 10^6$ and 10^5 cm^{-3} for OH and Cl , respectively) in such washing conditions, dependent on the amount of indoor illumination.

Practical Implications

Washing a floor with bleach leads to the introduction of a number of chlorinated gases to the room, including hypochlorous acid and chlorine. The chlorine levels of particles in the room also rise. The hypochlorous acid is observed to decay rapidly, implying uptake to room surfaces. Photochemical modeling indicates that significant levels of gas phase radicals may

form, dependent upon the amount of room light. Overall, bleach washing leads to enhanced oxidation of not only the washed surfaces but also of other surfaces in the room and of the air.

Introduction

Household bleach is largely a sodium hypochlorite (NaOCl) solution commonly formed by electrochemical oxidation of chloride in basic solutions. With a pKa of 7.4, hypochlorous acid (HOCl) dissociates to ClO^- in sufficiently basic solutions, i.e. as in commercial bleach solutions which have a pH significantly higher than the HOCl pKa. When chloride is present under sufficiently acidic or dry conditions, the aqueous-phase equilibrium between Cl_2 , HCl and HOCl/OCl^- shifts from HOCl/OCl^- to Cl_2 :



Please refer to Figure 1 for a detailed schematic of the reaction mechanisms prevalent in this work.

Active chlorine in the form of HOCl is a good oxidizing agent and is used widely as an antimicrobial agent in water systems, both for sanitizing drinking water and swimming pools. As well, it is used in household and commercial settings as a cleaning agent of both surfaces and clothes. HOCl exhibits a broad disinfecting ability given its reactivity with a number of types of biomolecules, leading to protein denaturation, radical production, and lipid peroxidation.^{1,2} Being a neutral species, it can more easily cross through cell membranes than its anionic counterpart, ClO^- . Specifically, HOCl is a good oxidizing agent of thiol groups leading to protein modification.³ It can also react with nitrogenated biomolecules, such as nitrogen-containing amino acids within DNA and RNA.^{3,4} Finally, HOCl is known to add across carbon-carbon double bonds forming chlorohydrins.² Thus, it can modify the properties of unsaturated lipid molecules, potentially affecting membrane properties and leading to oxidative stress and peroxidation via the formation of singlet molecular oxygen.^{5,6} HOCl is formed endogenously in the body by the myeloperoxidase enzyme present within classes of white blood cells.⁷ It is naturally released as an infection control agent given its anti-microbial properties.

From the photochemical perspective, small chlorinated molecules such as HOCl and Cl_2 are well known participants in atmospheric photochemical and multiphase reactions, such as

those that occur in the Antarctic Ozone Hole. Reactions of particular importance in that environment include the formation of Cl_2 from a heterogeneous reaction occurring on polar stratospheric cloud materials between HOCl and HCl :



This reaction proceeds extremely rapidly with uptake coefficients approaching unity on cold ice surfaces.⁸ The aqueous counterpart of reaction (2), i.e. reaction (1), is reasonably facile with a liquid-phase rate constant of $4 \times 10^4 \text{ M}^{-2}\text{s}^{-1}$ in the forward direction.² Once liberated, Cl_2 is sufficiently volatile that it is released to the gas phase where it can be readily photolyzed with near ultraviolet/visible light to form atomic chlorine that drives important ozone-destroying catalytic cycles. Chemistry of a similar nature also occurs in the tropospheric marine boundary layer.⁹

The role of chlorine-containing substances on the quality of indoor air has been investigated to only a small degree. One study has reported halogenated volatile organic compounds (VOCs) such as chloroform and carbon tetrachloride present in the headspace of bleach solutions and indoor settings, possibly arising from HOCl reacting with organic surfactant compounds added to bleach.^{10, 11} Similarly, it was found that chloroform can be released from the use of bleach in washing machines.¹² Additionally, emissions of chloroform have also been documented from other indoor water use activities, such as showering.^{13, 14} The general topic of the potential impacts arising from indoor cleaning agents has been discussed in a review which summarizes a number of reported incidents of inappropriate simultaneous use of bleach with ammoniated cleaners.¹⁵ This chemistry produces chloramines which are themselves disinfecting agents but can also have negative effects on human health. As well, there have been incidents in which acidity drives Reaction (1) forward, releasing chlorine gas.¹⁵ Finally, other chlorinated species, in particular chlorine dioxide, have been used for indoor disinfection¹⁶ with OClO exhibiting surface uptake capability.¹⁷

Given the small number of bleach-related indoor air studies, this paper addresses the impacts arising from the use of bleach as a surface-washing agent. We ask the following questions: What mixing ratios of reactive chlorinated gases such as Cl_2 and HOCl arise when bleach is used indoors? Is only the gas-phase composition of a room affected or are there also

impacts on the suspended particulate matter? Is there evidence for uptake of these gas-phase molecules by indoor surfaces? What photochemistry will result from the bleach emissions? What other species are released to the indoor environment when washing with bleach? Our goals are not to quantitatively assess the variability of emissions by surveying a wide range of bleach sources, or the variability of emissions that might arise from one bleach supplier. Instead, the focus is on identifying for the first time the suite of molecules that may be released to the indoor environment and to assess the behavior that they exhibit once released. One particular issue is the formation of oxidants, such as the OH and Cl radicals, that might arise through photolysis of chlorinated compounds. These radicals have the potential to increase the oxidizing capacity of indoor air and shorten the lifetime of VOCs.

Methods

1. Experimental

Experiments were conducted in a chemistry laboratory with continuous linoleum flooring, i.e. there are no seams or joints other than where the flooring meets the wall. The concrete ceiling and cinder block walls are painted with a gloss paint. The room has dimensions of 12.2 m long by 6.7 m wide by 3.8 m high, and contains scientific equipment as well as cabinets and benches. We estimate the free air volume is $292 \pm 15 \text{ m}^3$, where the uncertainty is associated largely with uncertainties in the volumes of the objects in the room. Lighting is provided by four large windows (each 2.9 m^2) and fluorescent lights. In the middle of the room, the lights levels were roughly 0.7 W/m^2 as measured with a spectral radiometer from 300 to 800 nm when the lights were on and the windows uncovered. Roughly half the flux came from the windows and half from the fluorescent lights. Building air is not recirculated. It enters the room through vents in the ceiling having first been chilled and filtered, and it mostly exits through two fume hoods in the room. The temperature in the room was typically between 23 and 26 °C and the relative humidity was less than 60%.

The air exchange rate was measured by allowing dry ice to sublime at a number of locations around the room until the CO₂ mixing ratio reached roughly 1000 ppm levels. Fans were deployed in the room to ensure homogeneous mixing of air. The remaining dry ice was then quickly removed from the room and the CO₂ level was monitored (Thermo Model 410i) as a function of time as it declined to background levels, without any human participants in the room.

Measurements were conducted four times in total across three different days. The CO₂ decay rates were highly reproducible across these replicates (Table 1), indicating minimal fluctuations in the building air exchange rate.

Experiments were conducted by washing the floor with a bleach solution prepared according to manufacturer recommendations, i.e. a 97% dilution of the stock solution using MilliQ water in a plastic bucket. Mopping (with a commercial mop) roughly half of the floor area ($37 \pm 5 \text{ m}^2$) was conducted rapidly, on the order of a minute or so. The height of the solution in the bucket was measured, yielding an average value of $200 \pm 44 \text{ mL}$ of solution applied in each experiment. Floor washing was conducted with at most three people in the room, and once the washing was complete the participants left the lab for roughly 30 to 45 minutes leaving the on-line instruments monitoring.

On-line measurements were conducted using instruments placed within the laboratory. A chemical ionization mass spectrometer (CIMS, Aerodyne Inc.), consisting of an iodide reagent ion source and a time-of-flight mass spectrometer (mass resolution of roughly 3000 (Th Th^{-1})) sampled room air without a sample line at a rate of 300 sccm and at a height of 1 m from the floor.¹⁸ This sample flow was diluted with 1700 sccm of nitrogen flow to ensure that the iodide reagent ion was always in excess (i.e. the average decrease in iodide reagent ion was $\sim 3\%$ during a typical mopping experiment). The iodide ion readily clusters with a large range of gas-phase species (X), including a variety of chlorinated compounds, detecting them as $\text{I}^-\cdot\text{X}$ adducts.^{19, 20} Positive identification of chlorinated gases is made using both high-resolution mass fitting of the mass spectra peaks and from the expected isotopic composition of the gases arising from the ³⁵Cl and ³⁷Cl isotopes.

Signals of Cl₂ were calibrated in a standard manner by supplying known amounts of gas from a glass bulb of known volume containing a dilute mixture of Cl₂ in nitrogen. The bulb was prepared by sequential dilution of a small quantity of pure Cl₂ using a capacitance monometer (MKS Inc.) mounted on a vacuum manifold to measure the dilution ratios.

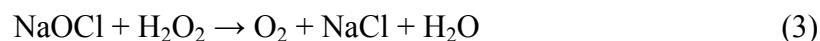
HOCl is not commercially available as a gas. As a result, HOCl mixing ratios were calibrated by converting HOCl to Cl₂ according to a modified procedure described previously.²¹ In particular, the HOCl and Cl₂ signals were observed when a 25 – 30 sccm nitrogen flow from a 1% by weight solution of NaOCl, buffered to pH 7.0 using a phosphate buffer, passes over a 10% by weight solution of NaCl containing 1% HCl and held at 0 °C. From the decrease in

HOCl signal and the associated increase in Cl₂ signal when passing over the chloride solution, the relative sensitivities of the CIMS to the two species can be determined. Using this information, as well as the Cl₂ instrumental sensitivity determined as described above, the HOCl sensitivity is assessed. We estimate absolute uncertainties in the reported Cl₂ mixing ratios to be on the order of $\pm 30\%$, arising from the uncertainties in the Cl₂ mixing ratio in the glass bulb and the calibration. The uncertainty in the HOCl mixing ratios is considerably higher than that for Cl₂, on the order of a factor two, i.e. $\pm 50\%$. In addition, the mixing ratios of HOCl reported in the paper are likely lower limits to the true values given that our calibration procedure assumes unity conversion of HOCl to Cl₂. Enhanced relative humidity had minor effects on the instrumental sensitivity, and the detection limits for Cl₂ and HOCl were 7 and 53 pptv respectively (at three sigma of the background).

Aerosol composition was monitored on-line by an Aerodyne aerosol mass spectrometer (AMS) with a compact time-of-flight mass spectrometer (C-ToF-AMS). Room air was sampled at a flow rate of 90 sccm with no sample line attached and with the inlet at 1.3 m from the floor. The AMS has been reviewed elsewhere.²² It operates using electron impact ionization of vaporized species that arise after a particle beam under vacuum impacts a heated vaporizer. The ions are then extracted into a time-of-flight mass spectrometer. The AMS is largely sensitive to non-refractory particles below one micron in diameter. The instrument is calibrated using known quantities of nitrate aerosol, and the mass loadings of other species are then assessed using relative ionization efficiencies which, for this work, are taken to be: 1.2 for sulfate, 1.4 for organics, and 1.3 for chlorine.²²

Gas-phase ozone (O₃) was measured using a Thermo monitor (Model 49i), and sub-micron aerosol size distributions were measured with a scanning mobility particle sizing (SMPS) system from TSI Inc. which included a model 3080 DMA and model 3025 CPC.

The hypochlorite composition of the commercial bleach solution was measured to be $8.7 \pm 0.1\%$ (by mass) by reacting the solution with excess H₂O₂ and measuring the volume of O₂ gas released:



2. Modeling

The modeling was carried out using the Indoor Chemical Model (INDCM) as described previously.^{23, 24} The zero-dimensional box-model uses a detailed chemical mechanism based on the Master Chemical Mechanism (MCM),^{25, 26} which is adapted for use indoors through the inclusion of terms that represent surface deposition, photolysis via attenuated light from outdoors and/or indoor lighting, exchange with outdoors and indoor emissions.

Although the MCM contains chemical degradation schemes for a number of halogenated hydrocarbons and considers reaction of the Cl radical with a range of alkanes, it does not contain comprehensive chlorine chemistry. Therefore, a recently developed module for studying chlorine chemistry at a coastal site in Hong Kong and designed for use in the MCM by Xue et al.²⁷ was added to the INDCM to address this gap. The new mechanism contains 205 reactions that represents the reactions of a range of VOCs with the Cl radical, as well as inorganic chemistry of both Cl and its precursors.

The model was modified to represent the dimensions of the laboratory, providing a surface area to volume ratio of $\sim 1.0 \text{ m}^{-1}$ to drive deposition of gases in the model. The observed air exchange rate of 12.74 h^{-1} was used and photolysis rates were calculated for indoor lighting according to the method described by Nazaroff and Cass.²⁸ The wavelength dependent quantum yields for the new chlorinated species were taken from the latest IUPAC evaluation.²⁹ Outdoor light was assumed to be attenuated by 97% in the UV and 90% in the visible when compared to outdoors,²³ though this assumption was tested as described later.

Outdoor VOC concentrations in the model were initialised according to Sarwar et al.,³⁰ whilst NO, NO₂, SO₂ and CO were taken from annual mean values from downtown or west Toronto³¹ and were set to 2.7, 13.5, 0.6 and 250 ppb respectively. Given the high air exchange rate, indoor steady-state NO and NO₂ mixing ratios are ~ 0.4 and 14 ppb respectively (at the high air exchange rate, indoor concentrations approximate those outdoors in the absence of indoor emissions, plus some NO is converted to NO₂ through reaction with O₃). Indoor ozone mixing ratios varied between 30 – 50 ppb over the course of the experiments and these measured mixing ratios were used to guide the model runs.

The new chlorine mechanism considers uptake of four species (NO₃, N₂O₅, ClONO₂ and HOCl) onto aerosol surfaces. The uptake coefficients were assumed to be 4×10^{-3} for NO₃ and 1×10^{-2} for the other three species.²⁷ During the experiments, SMPS data indicated the average value of aerosol surface area was $1.3 \times 10^{-4} \text{ cm}^2/\text{cm}^3$.

The amount of bleach solution used per mopping event and its concentration enabled an emission rate to be calculated. In particular, it was estimated that $\sim 7 \times 10^{-3}$ moles of HOCl were emitted into the room over 1 minute of mopping. Given the room volume, the emission rate of HOCl was calculated to be ~ 9.6 ppb/(cm³ s). Although the mopping lasted for only 1 minute, the emission into the room likely lasted longer, continuing as the bleach solution evaporates. The emission rate was therefore varied in the model to apply the relevant quantity of HOCl over time periods varying from 1-5 minutes. It was found that the shapes of the measured peaks were best reproduced when the emission was assumed to occur over 5 minutes. Therefore, the emission rate was set to ~ 1.92 ppb/cm³/s over 5 minutes.

As described below, Cl₂ was detected in the bleach solution headspace indicating it was present as an impurity. A direct emission of Cl₂ into the laboratory was therefore assumed. The sensitivity of the model was tested and the Cl₂ direct emission was found to be optimal at 5% of HOCl. Also, there are unlikely to be chemical or physical losses for Cl₂ within the room, given it appears (see below) to be removed at the rapid air exchange rate. We therefore assumed that deposition of Cl₂ to surfaces was negligible.

Experimental Results and Discussion

Figure 2A presents a typical mass spectrum obtained from the room air immediately after floor mopping, where mass spectral signals are normalized to reagent ion intensity and represent the difference in signal between when bleach is present and prior to its application. Figure 2B is a mass spectrum taken by passing a small flow of air through the bleach solution in a glass bubbler. Clearly identifiable peaks arise from iodide clustering with ClO, Cl₂, HOCl, ClNO₂, NHCl₂ and NCl₃. High mass resolution peak fitting has established the elemental composition of each ion (see SI Figure 1 and SI Table 1). Confirmation of molecular assignments comes from the chlorine isotopic abundances (see SI Table 2). We note that the largest peak in the spectrum aside from the reagent ion (not shown) is I·Cl which could arise from fragmentation of iodide clusters with chlorinated molecules. Although this needs to be confirmed, we believe that the iodide cluster with ClO may also arise from fragmentation of larger species, given that it is unlikely that a reactive radical would be measured at comparable signal intensities as other non-radical species, such as NHCl₂. However, we can not rule out the presence of ClO radical.

In Figure 3 temporal data are plotted from a typical set of mopping experiments. In Figure 3A, the data for Cl_2 and HOCl are represented by their reagent-ion-normalized signals and also by their mixing ratios in the room. As well, the normalized signal for ClO and Cl_2O (multiplied by 100) are included. In each case, the signal is that due to the strongest isotopic peak in the spectrum. Figure 3B shows the reagent-ion-normalized signals of the mixed chlorine-nitrogen species: ClNO_2 , NCl_3 and NHCl_2 .

At the start of each day of mopping experiments, a control experiment was conducted (at 11:19 am for the data in Figure 3), where the floor was mopped with MilliQ water and no bleach solution was applied. No large signals arise from this activity. Four subsequent bleach mopping experiments are shown in Figure 3, starting at roughly 11:50 AM, 12:30 PM, 1:15 PM and 2:00 PM. In each case, the chlorinated signals rise and then decay away as the air is flushed from the room.

Two of the most prominent peaks in the mopping experiment arise from Cl_2 and HOCl (Figure 2A). Interestingly, the HOCl signal is largely absent in the headspace spectrum (Figure 2B). These values are compared graphically in Figure 4 where we have referenced each maximum signal to that due to Cl_2 , for both the mopping and headspace data. It is seen that the mopping HOCl data are orders of magnitude higher than the headspace data, likely due to solution acidity and dilution effects. In particular, the pH of the diluted bleach solution was measured to be 11, i.e. HOCl will be fully dissociated into the ClO^- form and thus unlikely to volatilize. By contrast, we hypothesize that uptake of CO_2 from the air will acidify the solution after mopping, shifting the HOCl/ClO^- acid-base equilibrium over to the HOCl form which will be much more volatile (see Figure 1):



It is also possible that acidic species are present on the floor surfaces that affect the equilibrium. Evaporation of water from the drying bleach solution will also shift the equilibrium in Reaction (4) to the right. Finally, we note that the species Cl_2O was observed in both the mopping and headspace spectra with a maximum normalized signal of roughly 3×10^{-3} . Cl_2O is the anhydride of HOCl (see Figure 1):



While the presence of Cl_2 and HOCl (and Cl_2O) arising from a bleach solution is expected based on the chemistry presented in the Introduction, the large signals from a number of the other observed species is unexpected. ClNO_2 is a well known atmospheric constituent formed by a multiphase reaction between gaseous N_2O_5 and chloride-containing surfaces, such as sorbed HCl , or with NaCl (see Figure 1):³²



We note that ClNO_2 was also observed when we sampled the vapour directly from a bleach solution (see Figure 4) but, when accounting for experimental uncertainties and referencing to Cl_2 , the observed levels were not significantly higher after mopping. (Note that the comparison in Figure 4 is for reagent-ion-normalized signals allowing comparison between data sets measured under different conditions.) This leaves open the question that some ClNO_2 arose through reaction (6) occurring on surfaces in the room, where N_2O_5 would be formed by NO_3 reacting with NO_2 , NO_3 being generated by the reaction of outdoor NO_2 with O_3 . We note that this chemistry should occur readily in dark and dimly lit surroundings, such as indoors. ClNO_2 is not readily calibrated using instrumentation in our lab but we note that the normalized signal levels were similar to those for both HOCl and Cl_2 . This implies that the indoor mixing ratios of ClNO_2 could potentially significantly exceed those in the outdoor environment, that are normally in the sub-ppbv range.⁹

Also observed in both the room air after washing and in the bleach solution headspace were two chloramines, NHCl_2 and NCl_3 . These are known products from the reaction of HOCl with ammonia, as occurs in swimming pools that use chlorine as a disinfectant. Comparing the mopping and headspace conditions, the ratio of NCl_3 to NHCl_2 decreased in the mopping condition. It is difficult to establish why this ratio changes, perhaps arising from loss of HOCl through volatilization to the room after mopping (although loss of water would expect to push the equilibrium in the reverse direction) (see Figure 1):



In Figure 3, it is seen that all the species decay rapidly after the mopping is complete. The first-order decay rate constants of the different CIMS signals are presented in Table 1, along with the mean decay for CO₂ from independent experiments. The latter value is $(3.54 \pm 0.08) \times 10^{-3} \text{ s}^{-1}$ which is very rapid for an indoor setting but arises in a chemistry laboratory that requires fast air turnover. The decay constant is derived by fitting an exponential function to the decaying portion of the gas signal profile.

We note that within experimental uncertainties, the Cl₂ and NCl₃ signals decayed at the air exchange rate (see Figure 5, for Cl₂). This implies that on this timescale, these species do not undergo significant chemistry once liberated into the gas phase. In particular, the decay rates were the same regardless of whether the experiment was conducted with the lights on or off, or with the windows covered, indicating that there is no photochemical loss on the air exchange timescale. On the other hand, ClNO₂ and NHCl₂ decayed somewhat more slowly than did CO₂, perhaps indicative of slow desorption from surfaces in the lab. By contrast, the HOCl signals decayed at, on average, 1.4 times the room air exchange rate, again regardless of whether the room was illuminated or not. As well, the maximum mixing ratio reached by HOCl became progressively higher with subsequent moppings during the day (e.g. see Figure 3). We interpret this set of observations as indicative of increasing amounts of HOCl being liberated from the floor after each mopping event as well as, potentially, from different rates of multiphase chemistry occurring between gaseous HOCl and other surfaces present in the room. At this point we cannot determine whether the uptake from the gas phase leading to the rapid decay is via inorganic processes, e.g. HOCl reacting with surficial chloride, or with organic materials, or via redox reactions involving HOCl.³³ In either case, the increasing levels of HOCl in the room after subsequent floor washings likely arise because the HOCl is consuming reactive materials on the floor and/or other surfaces, leaving less available for subsequent reactions. Additionally, we note that the rate of HOCl surface chemistry occurs on the same order of magnitude as indoor ozone surface chemistry (typical decay rates of ozone due to removal by indoor surfaces range from $(0.22 - 2.0) \times 10^{-3} \text{ s}^{-1}$).³⁴

In the absence of surface reactions, the areas under the curves in Figure 3A are measures of the amount of Cl₂ and HOCl released to the room. Those integrated values are in Table 2 in units of ppbv minutes. A large amount of variability exists in the HOCl signal within a day (as

described above) and also between days. Cl_2 shows such variability between days as well, although not to the same degree as HOCl . This observation is deserving of additional study to systematically determine whether the changing nature of the surfaces being mopped was giving rise to different values in the gas phase. The active chlorine level of the undiluted bleach source used in this work was determined to be stable to the 0.2% (by mass) level over a period of 30 days, i.e. the variability in the amounts of HOCl released from one day to the next does not arise from changing strength of the bleach solution.

Figure 3C shows the aerosol particulate mass loadings observed in the room via aerosol mass spectrometry. As expected, there is no effect on the levels of organic and sulfate aerosol components that arise by mopping but, interestingly, the aerosol chlorine loading responded to each mopping experiment. As mentioned above, the amount of particulate chlorine mass plotted in Figure 3C is dependent on the relative ionization efficiency employed in the analysis and, for this reason, should be viewed with some caution given that we do know the form of the particulate chlorine. Aerosol chlorine may arise from either unreactive partitioning to the room particulates or else via reactive uptake of gas-phase chlorinated species. We note that there is a general increase in the levels of particulate chlorine with increasing total particulate mass loadings (see Figure SI-2) but this relationship could arise from either reactive or non-reactive partitioning processes. It is striking that the levels of particulate chlorine rise at comparable timescales to those of the gas-phase chlorine signals, indicating uptake on a timescale of minutes or less. They decay (see Table 1) at rates just slightly lower than those for CO_2 .

Implications for Indoor Chemistry

The implications of these observations for indoor chemistry are assessed using the modeling approach described above, with a focus on the data in Figure 3. When all the HOCl in the bleach solution was emitted into the room, the model predicted a gas-phase mixing ratio of HOCl of ~ 450 ppb, in excess of the measured values. Further, the predicted mixing ratios of HOCl did not increase with each application as shown by the measurements. As described above, this observation indicated that a significant amount of the HOCl was lost on the surface of the floor during mopping and also, that the efficiency of this surface loss decreased with time. Therefore, the model was run with varying amounts of HOCl assumed to stick to surfaces, with

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3 this proportion varying over time in line with the surface becoming saturated. The experimental
4 data were used to tune the model in this respect and the results are discussed below.

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7 To determine the optimal combination of model input parameters, each non-zero
8 experimental data point was paired with the equivalent model result for Cl_2 and HOCl and the
9 root mean square difference determined for all points in each peak. This value was then
10 normalised to the averaged experimental mixing ratio during each of the peaks so that
11 differences between experiment and model for the higher mixing ratio peaks didn't bias the
12 discrepancy between model and measurements. The results presented are those for which the
13 overall root mean square differences are minimised.

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16 The best results were found for the following conditions when it was assumed that the
17 amount of HOCl lost to the surface was 74%, 60%, 52% and 40% for peaks 1 – 4 respectively,
18 with a deposition velocity for HOCl to surfaces in the room of 0.035 cm s^{-1} (see Figure 6). This
19 latter value is very similar to that observed for O_3 deposition indoors.²⁴

20
21 The model can also be used to investigate the concentrations of radical species (OH , Cl
22 and ClO) formed in its chlorine mechanism. The photolysis of HOCl produces OH and Cl ,
23 leading to other radicals following reactions with VOCs (in the case of OH to produce HO_2 and
24 RO_2) and with O_3 (in the case of Cl to form ClO). Figure 7 shows the modeled concentrations of
25 OH , Cl and ClO . Note that in order to investigate the impact of light on the radical
26 concentrations, these figures are shown with four different lighting conditions:

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1. 3% and 10% of outdoor UV and visible light respectively plus indoor lighting
 2. 1.5% and 5% of outdoor UV and visible light respectively plus indoor lighting
 3. Just indoor lighting
 4. Dark

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60 The model runs illustrate that mopping events exert a prominent impact on the indoor
radical concentrations which vary significantly with the intensity and type of light. Radical
mixing ratios rise from their background levels immediately following the onset of mopping
events. With the highest intensity of light assumed (case 1), the OH radical concentration can
reach $2.0 \times 10^6 \text{ molec cm}^{-3}$ which is equivalent or higher than typical outdoor values, and very
much higher than typical indoor levels. Even in the absence of any outdoor light, mopping events

can lead to appreciable concentrations of radicals. Interestingly, mopping events in the dark can result in a slight decrease in the OH radical concentration.

We used the INDCM to investigate the controlling rates of reaction for HOCl and Cl₂ removal. For Cl₂, at least 98% is removed through air exchange with the remaining up to 2% via photolysis. This prediction is consistent with the data in Figure 5. For HOCl, around 88% is removed through air exchange with around 9% via dry deposition and 3% via removal on aerosol surfaces.

Finally, the loss rates of different VOCs via OH or Cl were compared to determine which oxidants are most important under these conditions. For isoprene, loss through reaction with OH was ~ 4 times faster than for Cl, ~ 2 times faster for propene and 1.3 times faster for CH₄. However, for *n*-butane, loss via reaction with Cl was ~ 3 times faster than loss via reaction with OH under experimental conditions.

Conclusions

Elevated levels of gas-phase Cl₂ and HOCl arise after floor washing with a commercial bleach solution. We note that the maximum mixing ratios reported were attained in a room with an extremely fast air exchange rate (12.7 h⁻¹), one to two orders of magnitude higher than that in other indoor environments, such as schools, offices and restaurants where bleach flooring washing may be employed. Proportionally higher mixing ratios than those observed will arise in indoor spaces with more commonly encountered air exchange rates. The elevated Cl₂ and HOCl levels lead to enhanced concentrations of reactive free radicals, OH, Cl and ClO, that will shorten the lifetimes of gas-phase organics with subsequent environmental effects, such as enhanced rates of secondary organic aerosol formation. The observations of elevated levels of gas-phase ClNO₂, NHCl₂ and NCl₃, as well as particulate chlorine, were unexpected. Beyond the scope of this work, it is important to study their formation routes and potential impacts.

Bleach is used commonly as an oxidative disinfectant. It is intriguing that its impacts appear not to be restricted to the surfaces to which the cleaning solution is applied. In addition, the uptake of HOCl onto other room surfaces has the potential to react with organics and amines present on those surfaces. We note that these potential surface reactions may be even more significant in a typical indoor environment where the surface-to-volume ratios are higher than the estimated surface-to-volume ratio for the room employed in the current study (~ 1.0 m⁻¹).

Subsequent Cl_2 and HOCl photochemistry will also significantly increase Cl and OH radical concentrations, enhancing the oxidizing capacity of the gas-phase. Overall, bleach washing leads to enhanced oxidation rates on both surfaces and in the gas phase.

Acknowledgements

Financial support from the Alfred P. Sloan Foundation is acknowledged. The help of S. Abbatt is acknowledged for measuring the active chlorine concentrations in the bleach solution.

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PROOF

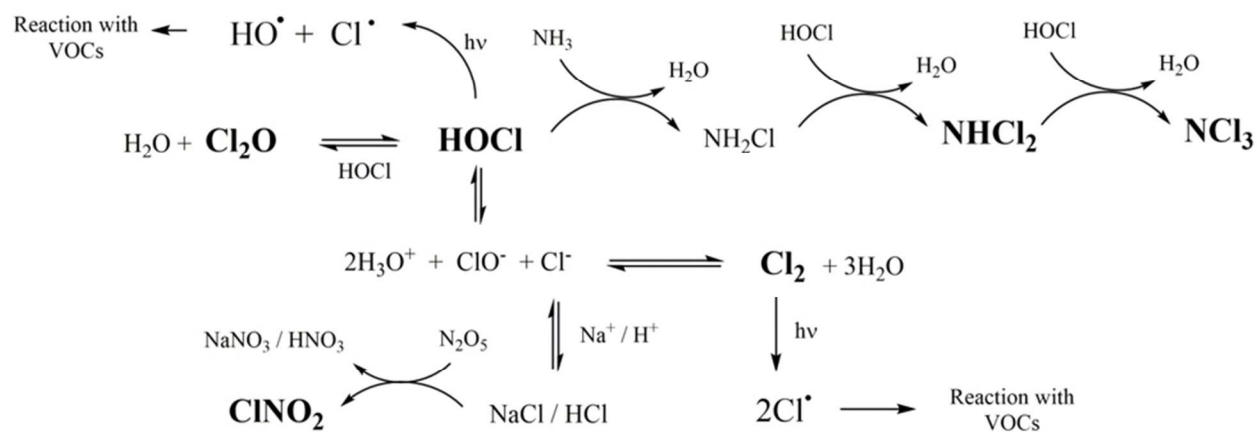


Figure 1. Detailed mechanism of indoor chemistry involving chlorine species. Species in bold were detected by the CIMS.

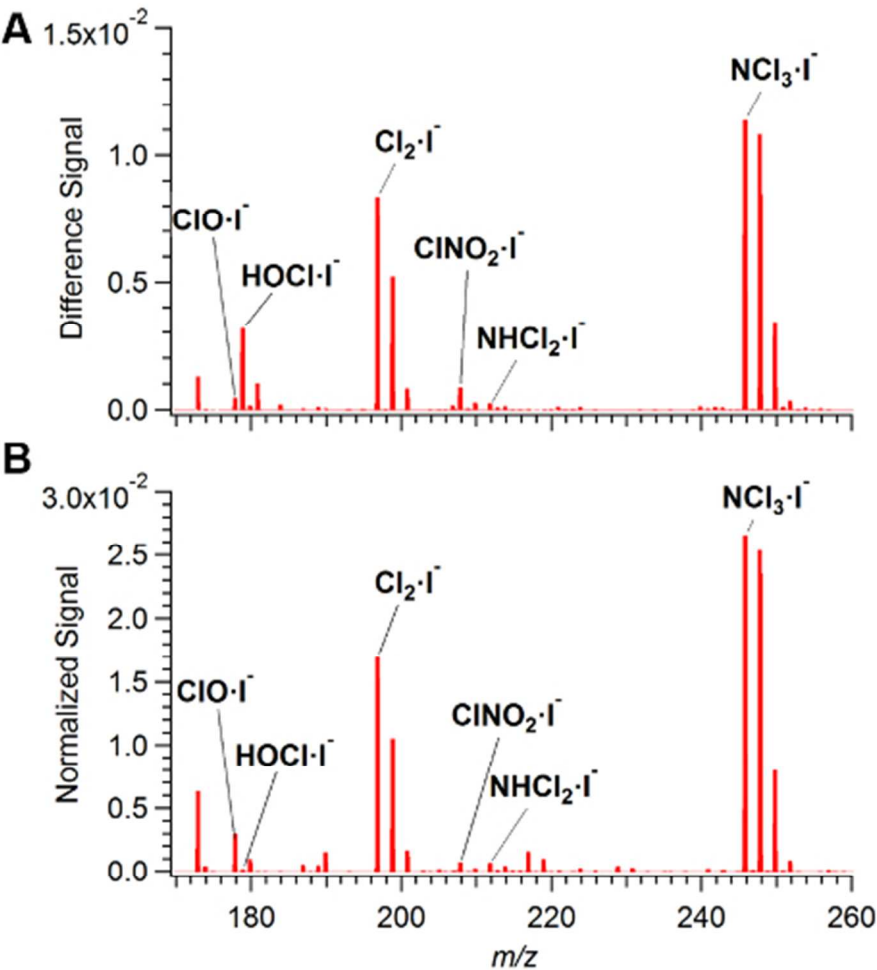


Figure 2. Typical iodide CIMS spectrum of (A) room air during a bleach mopping experiment and (B) the headspace from the bleach solution, using signals normalized by the reagent ion intensity. For (A), the signal plotted is the difference spectrum from before and after mopping, with positive signals indicative of an increase in signal.

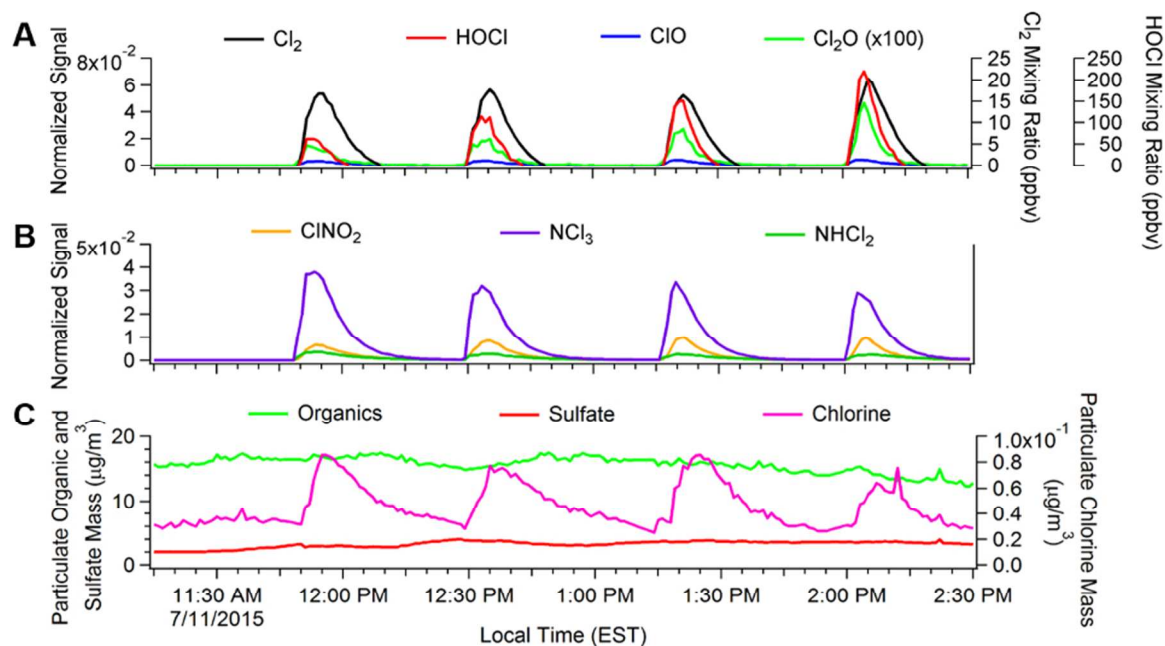


Figure 3. Time profiles of gaseous signals (A and B) and aerosol particulate signals (C) during a typical mopping experiment. Mopping was conducted with water once at the start of the experiment (11:19 AM) and with bleach four times subsequently. The mixing ratios of HOCl and Cl_2 are provided in Figure 3A. As discussed in the text, the particulate chlorine mass is dependent on the assumed value for relative ionization efficiency of chlorinated species in the AMS (here taken to be 1.3). Note that the Cl_2O trace in (A) was multiplied by 100.

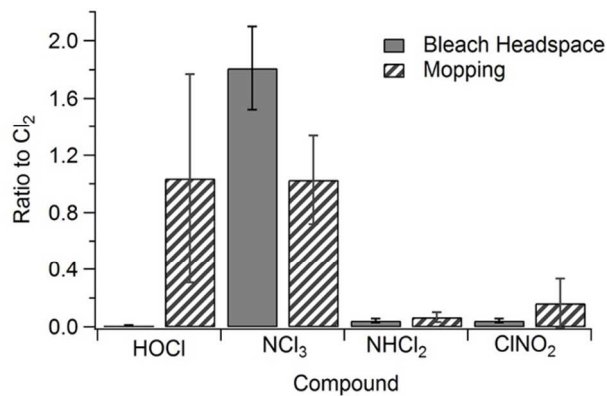


Figure 4. The average ratios of the signals of various species to Cl₂, both from the headspace and the mopping experiments. The error bars represent the variability ($\pm 1\sigma$) between multiple experiments.

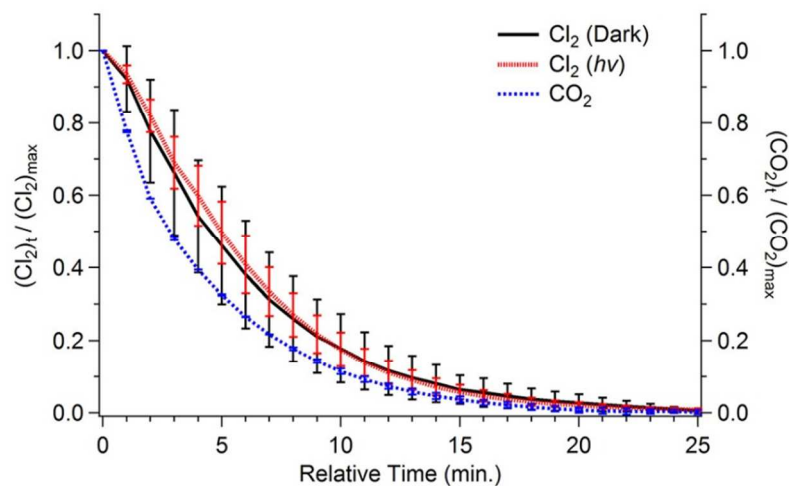


Figure 5. Plot of the Cl_2 signal normalized to its maximum value during a mopping experiment, as a function of time. Data are averaged for all experiments conducted and grouped by whether there was light present in the room or not. The error bars represent the variability ($\pm 1\sigma$) between multiple experiments.

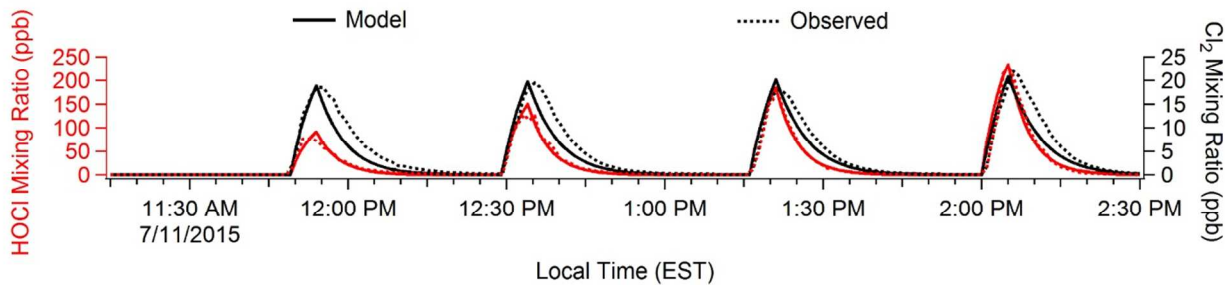
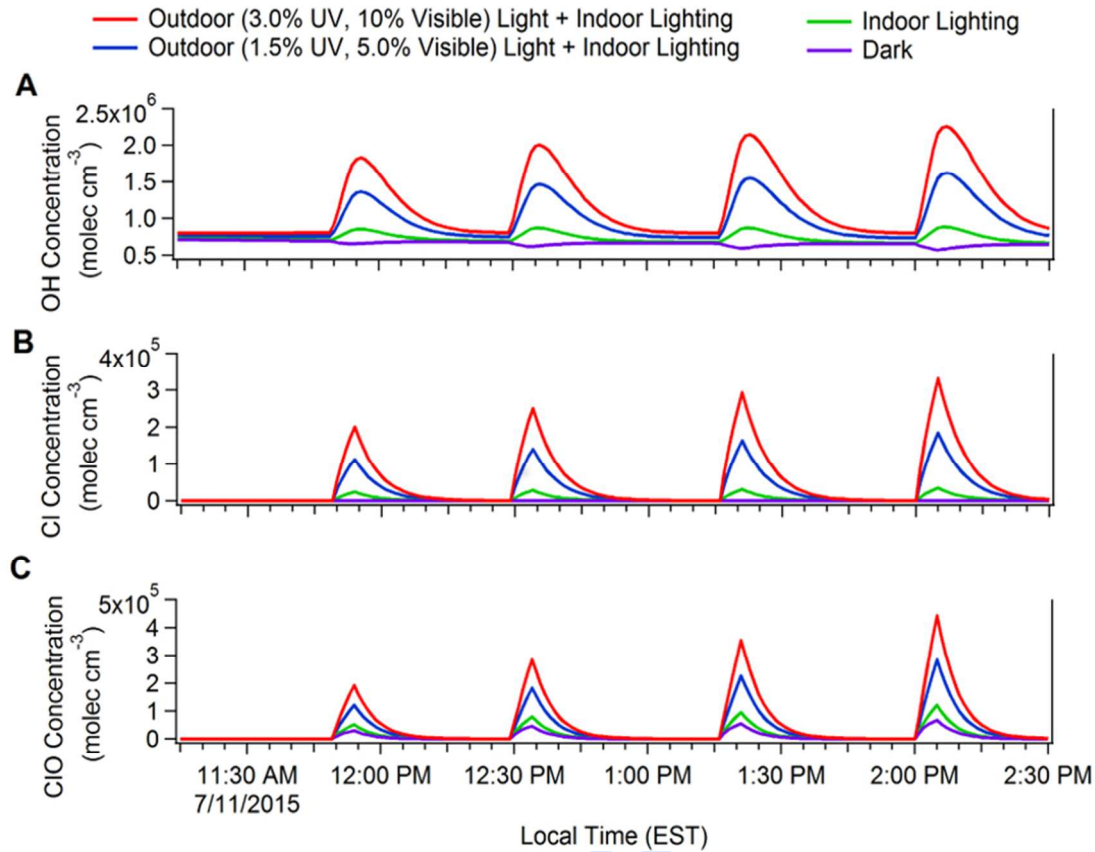


Figure 6. Plot of modeled and measured mixing ratios of HOCl and Cl₂ (see text for details).



32 Figure 7. Plot of modeled free radical concentrations under different light scenarios (see text for
33 details), for the model output presented in Figure 6.
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1 st Order Decay Rate Constants (10 ⁻³ , s ⁻¹)			
Compound	Dark (N = 4)	Room Light (N = 3)	Room Light + Sunlight (N = 5)
CO ₂	3.54 ± 0.08	--	--
Cl ₂	3.17 ± 0.90	2.99 ± 0.31	3.11 ± 0.72
HOCl	4.85 ± 1.86	5.11 ± 0.55	5.01 ± 1.21
ClO	4.75 ± 2.54	7.13 ± 0.67	5.40 ± 1.30
Cl ₂ O	2.75 ± 0.69	3.00 ± 0.24	2.72 ± 0.30
ClNO ₂	2.46 ± 0.37	2.35 ± 0.15	2.59 ± 0.62
NHCl ₂	2.04 ± 0.48	2.23 ± 1.07	1.83 ± 0.33
NCl ₃	3.37 ± 0.72	2.83 ± 1.33	3.12 ± 0.59
Particulate Chlorine	1.99 ± 0.88	2.27 ± 1.31	2.30 ± 0.32

Table 1. First-order decay rate constants for different species, after a mopping experiment. The uncertainties for each rate constant are ± 1σ, and the number of replicate experiments for each mopping experiment (N) is indicated.

Condition	Date of Expt	Cl ₂ (ppbv min)	HOCl (ppbv min)
Dark	7/4/2015 (#1)	230	1992
	7/4/2015 (#2)	246	2507
	7/11/2015 (#1)	211	697
	7/11/2015 (#2)	206	1073
Room Light Only	7/4/2015 (#3)	301	3666
	7/4/2015 (#4)	319	4318
	7/4/2015 (#5)	363	5255
Room Light + Ambient	6/16/2015 (#1)	445	2071
	6/16/2015 (#2)	461	3160
	6/16/2015 (#3)	422	3312
	7/11/2015 (#3)	191	1215
	7/11/2015 (#4)	221	1584

Table 2. Integrated areas of Cl₂ and HOCl signals, after each mopping experiment. The order in which each experiment was conducted on each day is indicated in brackets.

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Observations and impacts of bleach washing on indoor chlorine chemistry

J.P.S Wong,^a N. Carslaw,^b R. Zhao,^c S. Zhou,^d and J.P.D. Abbatt^{d,*}

*to whom correspondence should be addressed

^a School of Earth and Atmospheric Sciences, Georgia Institute of Technology, 311 Ferst Drive,
Atlanta, GA 30331 USA

^b Environment Department, University of York, Wentworth Way, Heslington, York YO10 5NG,
UK

^c Division of Chemistry and Chemical Engineering, California Institute of Technology, 1200 E
California Blvd, Pasadena, CA 91125 USA

^d Department of Chemistry, University of Toronto, 80 St. George Street, Toronto, ON M5S 3H6
Canada

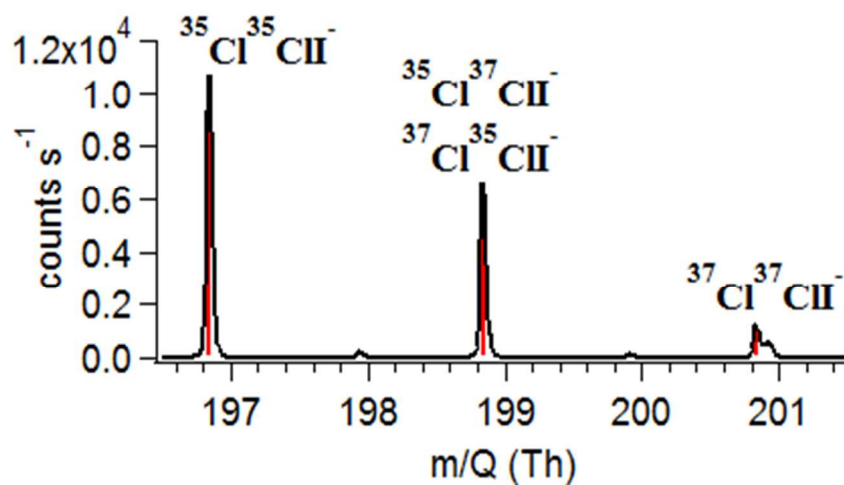


Figure SI-1. Expanded mass spectrum in the region of $\text{I}\cdot\text{Cl}_2^-$ signals, where the red line indicates the expected mass-to-charge ratio for the different isotopic combinations of Cl_2 .

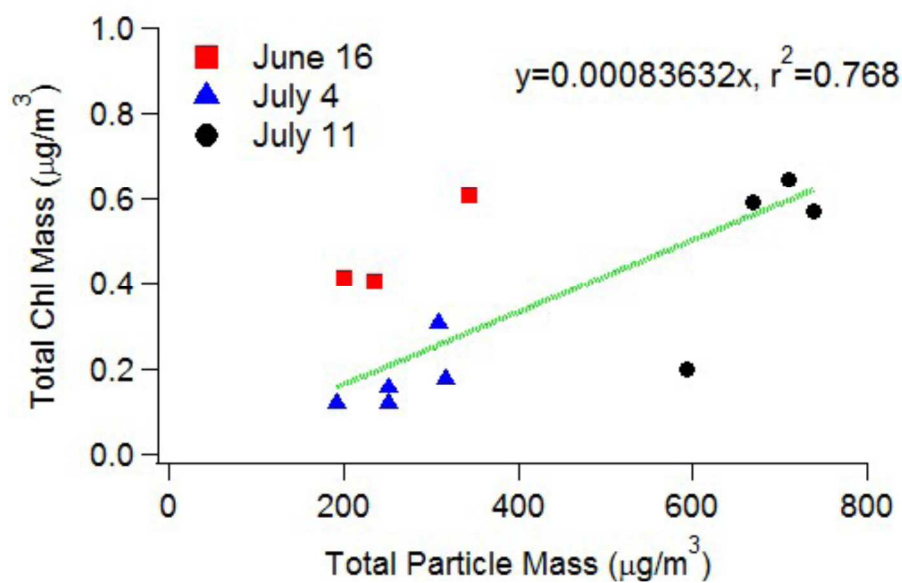


Figure SI-2. Plot of the total particulate chlorine mass as a function of total particle mass measured by the AMS.

Detected Formula	Expected m/Q	Observed m/Q
ClO ⁻	177.868789	177.8688
HOCl ⁻	178.876614	178.8766
Cl ₂ ⁻	196.842727	196.8427
ClNO ₂ ⁻	207.866778	207.8668
NHCl ₂ ⁻	211.853626	211.8536
NCl ₃ ⁻	245.814654	245.8147

SI Table 1. Expected and observed mass-to-charge ratios for the largest ion associated with different detected ions observed by the iodide CIMS.

Species	m/Q	Expected Abundance	Observed Abundance
³⁵ Cl ³⁵ Cl ⁻	196.84273	0.57373	0.572626
³⁵ Cl ³⁷ Cl ⁻ + ³⁷ Cl ³⁵ Cl ⁻	198.83923	0.367181	0.368879
³⁷ Cl ³⁷ Cl ⁻	200.83628	0.0587093	0.0584956

SI Table 2. Expected and observed abundances of ions observed by the iodide CIMS that are arising from the presence of Cl₂.